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11) Publication number:

0 218 410 A3

(12

EUROPEAN PATENT APPLICATION

(21) Application number: 86307302.9

(51) Int. Cl.3: A 61 K 31/44

(22) Date of filing: 23.09.86

- (30) Priority: 04.10.85 GB 8524508
- Date of publication of application: 15.04.87 Bulletin 87/16
- (8) Date of deferred publication of search report: 25.10.89
- Designated Contracting States:
 AT BE CH DE FR GB IT LI NL SE

- (1) Applicant: BEECHAM GROUP PLC Beecham House Great West Road Brentford Middlesex TW8 9BD(GB)
- (72) Inventor: Edwards, Peter John 7a Park Rise Leatherhead Surrey, KT22 7HZ(GB)
- (72) Inventor: Jeffryes, Carol Ann 14 Naseby Close IsleworthMiddlesex, TW7 4JQ(GB)
- 72) Inventor: Swain, Fiona Margaret 5 West Furlong KetteringNorthamptonshire, NN15 7LF(GB)
- (2) Representative: Russell, Brian John et al,
 Beecham Pharmaceuticals Great Burgh Yew Tree Bottom
 Road
 Epsom Surrey KT18 5XQ(GB)
- (54) Use of 1-hydroxy-2-pyridones in the treatment of acne.
- (5) A topical composition for application to skin affected by acne contains from 0.05 to 2% by weight of Octopirox together with a topically acceptable carrier. The composition is particularly useful for treating acne vulgaris.

EP 0 218 410 A3

EUROPEAN SEARCH REPORT

Application Number

EP 86 30 7302

	DOCUMENTS CONSI	DERED TO BE RELEVA	NT	
Category	Citation of document with i of relevant pa	ndication, where appropriate, ssages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. Cl. 4)
Χ	DE-A-3 140 954 (HC * Whole document *	ECHST)	1-7	A 61 K 31/44
Х	DIALOG INFORMATION De Haen Drug Data, 0141747, USAN Counc J. AM. MED. ASSOC.,	il: "Piroctone", &	1-7	
X	De Haen Drug Data,	il: "Piroctone", J.	1-7	
A	EP-A-0 117 080 (UN * Page 34, examples		1-7	-
A,D	FR-A-2 191 904 (HC * Page 12, lines 1- page 2, line 13 * &	20; page 1, line 1 -	1-7	
				TECHNICAL FIELDS SEARCHED (Inl. Cl.4)
				A 61 K
	The present search report has t	peen drawn up for all claims		
	Place of search	Date of completion of the search	1	Examuner
TH	E HAGUE	10-08-1989	GERI	LI P.F.M.
Y: pai do A: tec O: no	CATEGORY OF CITED DOCUMENTS X: particularly relevant if taken alone Y: particularly relevant if combined with another document of the same category A: technological background O: non-written disclosure P: intermediate document T: theory or principle underlying the invention E: earlier patent document, but published on, or after the filling date D: document cited in the application L: document cited for other reasons E: member of the same patent family, corresponding document		ished an, or	

EPO FORM (50) 03.62 (PO40)

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 Bulletin 87/16
- inventor: Edwards, Peter John, 7s Park Rise, Leetherhead Surrey, KT22 7HZ (GB) Inventor: Jeffryes, Carol Ann, 14 Naseby Close, isleworth§Middlesex, TW7 4JQ (GB) Inventor: Swain, Flons Margaret, 5 West Furlong, Kettering§Northamptonshire, NM15 7LF (GB)
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EP 0 218 410 A2

Composition

The present invention relates to a pharmaceutical composition for topical use, which contains a 1-hydroxy-2-pyridone or a salt thereof. In particular, the invention relates to a pharmaceutical composition for the treatment of acne.

US Patent No 4185106 discloses a class of 1-hydroxy2-pyridones which are described as being useful as anti-dandruff agents. It has now surprisingly been discovered that this class of materials is useful for the treatment of acne, which is nowhere mentioned or suggested in the aforementioned US Patent.

Accordingly, the present invention provides a topical composition suitable for application to skin which is affected by acne, comprising from 0.05 to 2% by weight of a compound of formula (I).

$$\begin{array}{c}
R_2 \\
R_1 \\
OH
\end{array}$$

$$\begin{array}{c}
R_4 \\
OH
\end{array}$$

or a topically acceptable salt thereof in which R₁ is hydrogen, alkyl of 1 to 17 carbon atoms, alkenyl of 2 to 17 carbon atoms, cycloakyl of 5 to 8 carbon atoms, bicycloalkyl of 7 to 9 carbon atoms, cycloalkylalkyl of 1 to 4 alkyl carbon atoms, the cycloalkyl groups being optionally substituted by alkyl groups of 1 to 4 carbon atoms, aryl, aralkyl of 1 to 4 alkyl carbon atoms, arylalkenyl of 2 to 4 alkenyl carbon atoms, arylakyl or arylthio-alkyl of 1 to 4 alkyl carbon atoms, benzhydryl, phenylsulfonylalkyl of 1 to 4 alkyl carbon atoms, benzhydryl, phenylsulfonylalkyl of 1 to 4 alkyl carbon atoms, furyl or furylalkenyl of 2 to 4 alkenyl carbon atoms, all the aryl groups mentioned being optionally substituted by alkyl of 1 to 4 carbon atoms, by alkoxy of 1 to 4 carbon atoms, by nitro, cyano or halogen;

 R_2 is hydrogen, alkyl of 1 to 4 carbon atoms, alkenyl or alkinyl of 2 to 4 carbon atoms, halogen or benzyl; R_3 is hydrogen, alkyl of 1 to 4 carbon atoms or phenyl; and

R₄ is hydrogen, alkyl of 1 to 4 carbon atoms, alkenyl of 2 to 4 carbon atoms, methoxymethyl, halogen or benzyl,

together with a topically acceptable carrier.

Preferred and exemplified compounds of formula (I) are those which are disclosed in the aforementioned US Patent No 4185106.

A particularly preferred compound of formula (I) is 1-hydroxy-4-methyl-6-(2,4,4-trimethyl pentyl)2(IH)-pyridone ethanolamine salt.

The preferred quantity of the compound of formula (I) or salt thereof in the composition of the invention is from 0.05 to 0.5% by weight, more preferably from 0.2 to 0.5% by weight.

In a further aspect of the invention, there is provided the use of a compound of formula (I), as hereinbefore defined, for the manufacture of a pharmaceutical composition for treating acne in humans, preferably acne in which the organism Propionibacterium acnes is implicated.

In a still further aspect of the invention, there is provided a method of treating acne in humans comprising applying a topical composition containing a compound of formula (I)or a salt thereof to the skin of a human suffering from acne.

A particularly preferred use for the composition of the invention is for the treatment of acne vulgaris, which is a polymorphic skin eruption characterised clinically by blackheads, white heads, papules, nodules, cysts and scars occuring particularly on areas of the skin rich in sebaceous glands, such as the face, forehead and back.

The topical composition of the invention may be presented in a wide variety of different forms, for example, creams, gels, ointments, lotions, sticks, soaps (liquid or solid), bath additives, shower gels, cleansing pads, impregnated wipes, face packs, shaving foams, aftershaves, atomiser sprays and other conventional cosmetic formulations.

The major requirement in the composition of the invention is that the topically acceptable carrier (which can be any ingredient conventionally used in the abovementioned compositions) should be non-irritant to an acne sufferer.

Normally, the composition of the invention would be applied two or perhaps three times daily, in accordance with conventional application techniques for topical formulations. The dosage level of active ingredient will depend primarily on whether the composition is a 'leave on' material, such as an

ointment, or a 'rinse-off' material, such as a soap.

Generally speaking, the dose for a 'rinse-off'

formulation would be two or three times that of a

'leave-on' formulation.

Compositions of the invention may be produced by conventional techniques for the manufacture of pharmaceuticals or cosmetics, usually involving admixture of the various ingredients to obtain a uniform composition.

The invention is now illustrated by the following Examples:

Example 1

Ge l	w/w
	per cent
¹ Octopirox	0.25
Menthol	10.00
DEA-oleth-3 phosphate	2.50
Hydroxymronylcellulose	2.50
2Hydroxypropylcellulose Amphoteric - 1	5.00
Water	39.75
Ethanol (96%)	40.00

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Ex	am	ÞΤ	е	

	w/w
Cream	per cent
Laneth - 10 Lanolin alcohol Cetvl alcohol Polawax Myristyl myristate Octopirox Resorcinol mono-acetate Magnesium aluminium silicate Methyl paraben Sulphur Perfume Water	2.00 0.50 5.50 6.00 2.00 0.25 0.2 4.00 0.20 1.40 q.s.
····	

Preparation: Dissolve the Octopirox in the propylene glycol and then add the rest of the oil phase ingredients. Add the magnesium aluminium silicate to the water at 75°C and disperse under shear again to dispense. Combine the phases and emulsify at 70°C, adding the perfume at 50°C.

- 1 Trade Mark of Hoescht for 1-hydroxy-4-methyl-6-(2,4,4-trimethyl pentyl) 2(1H)-pyridone ethanolamine salt.
- 2 Amphoteric-1 is the CTFA adopted name for cocoamphoglycinate.
- 3 Laneth-10 is the CTFA adopted name for glyceryl lanolate.
- 4 Polawax is a Trade Mark of Croda Chemicals Ltd.

Example 3

		W/W
<u>A</u>	erosol shaving cream	per cent
Part A	Stearic acid Lauric acid Liquid lanolin	4.0 2.0 1.0
	(1Cromeen Triethanolamine Octopirox Water (deionized) Perfume	3.0 2.5 0.5 87.0 q.s.
	Concentrate	92.0
	² Propellents 12/114 (40:60)	8.0

1 Cromeen (Croda Chemicals Ltd) is a substituted

alkyl amine derivative of various lanolin acids.

Example 4

Hydrocarbon-propelled aerosol shaving foam

		w/w
		per cent
D+	a Chalmitic acid	5.0
Part	A {Palmitic acid {Lauric acid	1.0
D h	B Sodium lauryl sulphate Polyethylene glycol (400) monolaurate Polyacrylic acid (40% aq) mol. wt 100 0	1.0
Part	b Sodium radific Surphiese (400) monolaurate	0.5
	polyethylene glycol (40% ag) mol. wt 100 0	00 1.5
	Polyacrylic acid (40% aq) mozi we asset	2.0
	Triethanolamine	0.8
	Potassium hydroxide	5.0
	Glycerol Octopirox Water (deionized)	0.5
	Octopirox	2.8
	(Water (deionized)	
	\Perfume .	q.s.
	Concentrate	96.9
	Propellants, isobutane/propane	3.1

Propellent 12 - Dichlorodifluoromethane. (B.P.).
Propellent 114- Dichlorotetrafluoromethane. (B.P.)

<u>Preparation:</u> Heat parts A and B separately to 75°C. Add A to B with vigorous stirring and allow to cool to 35°C, when the perfume is added. The aerosol container is charged when the concentrate has reached room temperature.

Example 5

	W/W
After shave lotion	per cent
Octopirox	0.25
Ethyl alcohol, specially denatured	60
Propylene glycol	3
Water, demineralised	35.75
Perfume	1

Preparation: Dissolve the perfume and propylene glycol in the alcohol and add the water slowly, stirring well to avoid locally high concentrations of water precipitating the less soluble components of the perfume. Allow the solution to stand for several hours at about 4°C, then filter.

Example 6

	w/w
Bath Liquid	per cent
Octopirox Sodium lauryl ether sulphate (28% Coconut diethanolamide	3
Perfume Citric acid Colour, preservative, emollients,	1-2 q.s. to pH 7
Sodium chloride	q.s. q.s. to required viscosity
Water	to 100

Example 7	w/w
Lotion	Per cent
Octopirox Alcohol Aluminium chlorhydroxyallantoinate Propylene glycol, Menthol Aluminium chlorhydrate (50%) Hydroxypropylmethylcellulose (3%) Mica (and) titanium dioxide Pefume, colour, preservative	0.25 43.00 0.20 3.00 0.05 5.00 47.75 1.00 q.s.

Example 8

	<u>w/w</u>
Stick	per cent
Sodium stearate Ethyl alcohol Propylene glycol Isopropyl myristate Octopirox Perfume	8.00 74.75 10.00 5.00 0.25 2.00

Procedure: Slurry the soap in the cold with organic solvents and Octopirox and then heat to 60° - 75°C Stir the mass while hot until clear. Add fragrance and colour as desired at 5° - 8°C above the set point of the stick. When it is uniform, pour the soap solution into moulds and allow to cool. Sodium stearate can be prepared in situ but critical control is required to avoid excess alkali or fatty acid.

Example 9	•		<u>w/w</u>
			per cent
Aerosol			. 0.25
0-1	•	•	0.25
Octopirox			2.00
Propylene glycol			57.25
Alcohol (99% V/V)	1		0.50
Perfume			40.00
Propellant 12			•••

Example 10	<u>w/w</u>
Clear gel face mask	per cent
Sodium magnesium sililcate PEG - 75 Octopirox	8.00 1.00 0.20 5.00
Alcohol Carbomer	to pH 7.5 to 100
Water perfume, colour, preservative	q.s.

Anti-microbial activity

To demonstrate the effectiveness of the preferred compound, Octopirox, of the composition of the present invention, the compound was subjected to <u>in vitro</u> evaluation by agar diffusion against <u>P. acnes</u> and S. aureus.

Method

Octopirox was evaluated at the 0.2%w/v level in either 10% ethanol or 10% *Tween 20.

O.1 ml of each solution was placed in a 1 cm diameter well in Brain Heart Infusion Agar (OXOID) seeded with either Propionibacterium acnes (strain 737) or Staphylococcus aureus (NCTC 6738).

*Tween is a trade mark of Atlas; Tween 20 is polyoxyethylene sorbitan monolaurate.

The plates containing Staph. aureus were incubated aerobically for 24 hours at 37°C and those seeded with P. acnes anaerobically for 48 hours at 37°C.

Results

Zone of Inhibition diameter (mm) (N=2a)

	P.acnes		S.aureu	<u>s</u> 10% Tween
	10% IMS	10% Tween	104 1W2	104 IMEE!!
No antimicrobial	NZ	NZ*	NZ	NZ
Octopirox	20.6	30	19	22.7

NZ = No zone of inhibition

* Zone of precipitation resulting from extracellular esterase activity.

Conclusion

The results demonstrate that Octopirox is effective against the organism <u>P.acnes</u> which is associated with the occurence of acne in humans.

Activity of Octopirox VS P. Acnes in the presence of an artificial sebum composition

Method

0.1ml of the test solutions/suspensions listed below were incorporated into 1cm wells cut into the surface of 245 x 245cm assay plates of brain heart infusion agar seeded with P.Acnes (strain 737) at a level of approx 10 6 cfu/ml. Zone of inhibition diameters were assessed after 48 hours anaerobic incubation at 37°C.

Test Agents

- 1. Octopirox (0.2%w/v) in 20% ethanolic solution.
- 2. As 1 above but also containing 10% artificial sebum.
- 3. Control 20% ethanol.
- 4. Control 20% ethanol + 10% artificial sebum.

Results

	mean zone diameter(mm)(n=		
AGENT	-sebum	+10% sebum	
Octopirox (0.2%)	18.2	18.7	
20% ethanol	No zone	No zone	
20% ethanol + 10% Artificial sebum	No zone	No zone	

Conclusion The results clearly demonstrate the ability of Octopirox to retain activity against P. Acnes in the presence of an artificial sebum composition.

The artificial sebum used in the above test method has the following composition:

Ingredient	8 W/W
Triglyceride Mix (1)	36
Fatty Acid Mix (2)	24
Cholesterol	4
Lanolin	8
Squalene	12
Glycerol	8
Water	to 100%

Triglyceride Mix (1)

Glycerol	palmitate	10	g
Glycerol	oleate	10	g

Fatty Acid Mix (2)

Palmitic Acid	10	g
Oleic Acid	5	g
Myristic Acid	5	g

 A topical composition suitable for application to skin which is affected by acne, comprising from 0.05 to 2% by weight of a compound of formula (I).

$$\begin{array}{c}
R_2 \\
R_1 \\
0
\end{array}$$

$$\begin{array}{c}
R_4 \\
0
\end{array}$$

or a topically acceptable salt thereof in which R₁ is hydrogen, alkyl of 1 to 17 carbon atoms, alkenyl of 2 to 17 carbon atoms, cycloakyl of 5 to 8 carbon atoms, bicycloalkyl of 7 to 9 carbon atoms, cycloalkylalkyl of 1 to 4 alkyl carbon atoms, the cycloalkyl groups being optionally substituted by alkyl groups of 1 to 4 carbon atoms, aryl, aralkyl of 1 to 4 alkyl carbon atoms, arylalkenyl of 2 to 4 alkenyl carbon atoms, aryloxy-alkyl or arylthio-alkyl of 1 to 4 alkyl carbon atoms, benzhydryl, phenylsulfonylalkyl of 1 to 4 alkyl carbon atoms, furyl or furylalkenyl of 2 to 4 alkenyl carbon atoms, all the aryl groups mentioned being optionally substituted by alkyl of 1 to 4 carbon atoms, by alkoxy of 1 to 4 carbon atoms, by nitro, cyano or halogen;

R₂ is hydrogen, alkyl of 1 to 4 carbon atoms, alkenyl or alkinyl of 2 to 4 carbon atoms, halogen or benzyl;
R₃ is hydrogen, alkyl of 1 to 4 carbon atoms or phenyl; and

R₄ is hydrogen, alkyl of 1 to 4 carbon atoms, alkenyl of 2 to 4 carbon atoms, methoxymethyl, halogen or benzyl,

together with a topically acceptable carrier.

- 2. A composition according to claim 1, in which the compound of formula (I) is 1-hydroxy-4-methyl-6-(2,4,4,-trimethyl pentyl)2(IH)-pyridone ethanolamine salt.
- 3. A composition according to claim 1 or 2, in which the compound of formula (I) or salt thereof is present in an amount of from 0.05 to 0.5% by weight.
- 4. A composition according to any one of claims 1 to 3 in the form of a cream, gel, ointment or lotion.
- 5. The use of a compound of formula (I) or salt thereof, as defined in claim 1, for the manufacture of a pharmaceutical composition for treating acne in humans.
- 6. The use according to claim 5, in which the organism implicated in acne is <u>Propionibacterium</u> acnes.
- 7. The use according to claim 5, in which the composition is for the treatment of acne vulgaris.